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i.	Title of the invention	COATING	
1	Name of your agent (gryous bave one)	Andrew Steven BOOTH	
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COATING

The present invention relates to a method of forming a coating on a substrate. More specifically but not exclusively, the invention relates to a method of forming a corrosion resistant coating on a machined part used, for example, in a vacuum pump.

Vacuum pumps are used in the manufacture of semiconductor chips to facilitate the control of the various environments that the chip must be exposed to during manufacture. Such pumps are typically manufactured using cast from and steel components, many of which are precision engineered to ensure optimum performance of the pump. Plastic based parts may also be used as components in vacuum pumps under certain conditions as described below.

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Iron castings and steels have for a long time been used in the manufacture of component parts for equipment used in a wide range of industries, including the petro-chemical and semiconductor industries. These parts are cheap, exhibit good thermal and thermo-mechanical properties and are relatively easy to form. However, in the semiconductor industries the increasing use of high flow rates of process gases (such as chlorine, boron-trichloride, hydrogen bromide, fluorine and chlorine-trifluoride) together with the associated elevated temperatures and pressures required have resulted in the severe corrosion of the Iron and steel component parts. Such corrosion leads to equipment failure, leakage of process chemicals and possible process contamination, and reduced process efficiency, as well as the costs associated with un-planned downtime.

In an attempt to minimise these problems, it has been common practice within many industries to passively protect many of the component parts, since this represents a cheaper alternative to the more expensive active protection that is available. The use of an aluminium coating on iron castings and steels, for example, has been used in a variety of industries to provide good corrosion and heat resistance. In addition, hot-sprayed ceramic coatings applied directly to the

metal surface have also been used to protect Iron and steel castings in abrasive and high temperature applications.

It has also been suggested that corrosion problems can be overcome by substituting the iron and steel parts with more expensive materials such as nickel rich iron base alloys, Monel, Inconel or higher nickel content alloys. However, these materials are expensive and do not represent a cost efficient alternative for use as component parts.

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More recently there has been a move towards the use of plastics-based component parts in a variety of industries in an attempt to replace the metal component parts traditionally used. The versatile nature of plastics means that they can be used to replace metal parts for a variety of reasons. Plastic parts can be manufactured by a variety of means and can be tailored to meet a number of application requirements. In addition their reduced weight and cost in comparison to metals means that they represent an attractive alternative in the manufacture of machine parts. However, because of the susceptibility of these materials to the intensively corrosive, oxidative and aggressive environments encountered in the semi-conductor industry, their use in equipment in this industry has been limited. Most plastic materials will readily wear in the presence of abrasive particles and many hydrocarbon-based plastics may spontaneously combust in the presence of fluorine or oxygen gas.

Many attempts have been made to impart wear and corrosion resistance to a number of plastics materials, the provision of ceramic coatings being particularly popular. However, the application of ceramic coatings to plastic substrates has not always proved easy because, unlike metal surfaces, it is difficult to form ceramic coatings on plastic surfaces that exhibit good adherence and do not flake off in service. This is thought to be due to the non-conductive nature of the plastic surface, which results in the build up of electrostatic charge during the spraying process and acts to repel the sprayed ceramic particles.

There is therefore a need for a corrosion resistant coating that can be easily applied to a metal or plastic substrate and which exhibits good adhesion thereto.

In one aspect, the present invention provides a method of forming a coating on a metallic or plastics substrate, the method comprising the steps of applying a metallic layer to the substrate and forming the coating from the metallic layer by subjecting the metallic layer to electrolytic plasma oxidation.

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The present invention thus provides a simple and convenient technique for forming an anti-corresive coating on a metallic or plastics compensation a vacuum pump. By the term "anti-corresive" it should be understood to mean that the coating to capable of withstanding wear and degradation as a result of exposure to abrasive particles and gases such as fluorine, chlorine-trifluoride, tungsten-hexatiuoride, chlorine, boron-trichloride, hydrogen bromide, oxygen and the like. The coating can be conveniently formed from any sultable transcriblayer forming metal or alloys thereof. By the term "barrier layer-forming metal" it should be understood to mean those metals and their alloys (such so Ai, Mg, Ti, Ta, Li, Nb, Hi, Sb, W, Mc, V, Si), the surfaces of which naturally react with elements of the environment in which they are placed (such as oxygen) to form a coating layer, which further inhibits the reaction of the metal surface with said reactive environmental elements.

The technique of electrolytic plasma oxidation (EPO) is known by various other names, for example anodic-plasma oxidation (APO), anodic spark oxidation (ASO), micro-arc oxidation (MAO). During this technique, a partial oxygen plasma forms at the metal/gas/electrolyte phase boundary and results in the creation of a ceramic oxide layer. The metal ion in the ceramic oxide layer is derived from the metal and the oxygen formed during the anodic reaction of the aqueous electrolyte at the metal surface. At temperatures of 7000K associated with the formation of the plasma, the ceramic oxide exists in a molten state. This means that the molten ceramic oxide can achieve intimate contact with the metal surface at the metal/oxide boundary, which means that the molten ceramic oxide has sufficient time to contract and form a sintered ceramic oxide layer with few pores. At the

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electrolyte/oxide boundary, however, the moiten ceramic oxide is quickly cooled by the electrolyte and the gases flowing away, notably oxygen and water vapour, leaving an oxide ceramic layer with increased porosity.

Thus, the ceramic oxide coating so formed is itself characterised by three layers or regions. The first is a transitional layer between the metallic layer and the coating where the metal surface has been transformed, resulting in excellent adhesion for the coating. The second is the functional layer, comprising a sintered ceramic oxide containing hard crystallites that give the coating its high hardness and wear redistance characteristics. The third is the curiage happy, which has lower hardness and higher porceity than the functional layer

It will be appreciated from the foregoing that the paramic oxide coating is atomically bound to the underlying metallic layer and is formed from the surface of the installic layer. This means that the commit oxide coating so produced exhibits greater adhesion to the underlying metallic layer than would be formed from externally applied sprayed ceramic coating. The periodic pales parallel superior surface properties such as extreme hardness, very low wear, detonation and cavitation resistance, good corrosion and heat resistance, high dielectric strength and a low coefficient of friction. In addition, it is also resistant to corrosion from halogens, inter-halogen compounds and other semiconductor processing chemicals excited by plasma.

From the foregoing it will be appreciated that the external surface of the coating is in some applications characterised by a low porosity. In such situations outgassing from the coated substrate material is minimised. In other applications, the external surface of the coating may be irregular and exhibit some porosity. In order to ensure extreme hardness, low wear and good corrosion resistance, the external surface of this coating may be removed by grinding to expose the underlying sintered ceramic oxide layer, which provides the superior surface properties referred to above.

Alternatively, where the external surface of the coating exhibits some porosity it can serve as a matrix for application of an optional layer of a composite nature. In such situations, materials suitable for forming the composite layer include a lubricant or paint, for example. It will be appreciated that the pore sizes of the external surface of the second layer are of a size that are capable of retaining the material of the third layer. Other examples of such composite coatings include lubricants such as fluorocarbons, polytetrafluoroethylene (PTFE), molybdanum disulfide (MoS₂), graphite and the like, which are retained by the porous external surface of the coating. The optional layer is preferably formed directly over the docating, the occasing providing a key for the adirection of this additional layer.

The metallic layer is suitably applied by depositing a layer of the barrier layerforming metal or alloy thereof directly or indirectly (depending on substrate) onto the substrate surface to a thickness of preferably less than 100 µm. The matellic layer is preferably deposited unito the surface of the outstrate using one of (1), (sifting or compression of metallic powder or wrapping of the foil onto a liquid dulingsive, after it has been applied to the surface (ii), electrolytic deposition onto an initially deposited metal layer (iii), spraying techniques such as sputtering, plaama-spraying, arc-spraying, flame-spraying, vacuum-metallising, ion-vapour deposition, high velocity oxyfuel-spraying, cold gas-spray; combinations thereof and the like, which are well known to a skilled person. These methods ensure that the metal or alloy thereof is both well adhered to and does not degrade the underlying substrate. Whatever procedure or combination thereof adopted, the parameters must be adjusted to values suitable to obtain homogeneous coatings. with low porosity value and free of cast-in (embedded) particles, oxides and cracks that will compromise the formation of the ceramic oxide coating by electrolytic plasma oxidation. For both metal and plastic substrates, the deposition of a metallic layer on the surface of the substrate has little effect on the bulk temperature of the substrate, thereby preventing distortion thereof. employing the hot spraying techniques, the superior wetting properties of the molten metal particles on the substrate surface, when compared to conventionally

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sprayed ceramic particles, lead to the formation of a metallic layer having a low porosity.

As indicated above, the coating is formed by electrolytic plasma oxidation of the surface of the metallic layer. The coating is suitably formed by immersing an anodically charged metal coated part in an alkaline electrolyte (e.g., aqueous solution of an alkali metal hydroxide and sodium silicate) using a stainless steel V625 to eacone in egation OA in acting and epotential representations are partially as the country electrons and applying an AC voltage in excess of 250V During this technique, a partial oxygen plasma forms at the modeliges/stackulyta places boundary and results in the medium of a commit oxide layer. The metal ion in the ceramic oxide layer is derived from the metal and the oxygen formed during the anodic reaction of the aqueous electrolyts at the metal surface. At temperatures of 7000K associated with the formation of the plasma, the ceramic oxide exists in a molten state. This means that the molton auranio pidde con achievo intimete contact with the instal euriado at 1 🔫 metal/oxide boundary, which means that the molten ceramic oxide has sufficient. time to contract and form a cintered peremie cride layer with few person. At the electrolyte/oxide boundary, however, the molten caramic axide is quickly cuoled by the electrolyte and the gases flowing away, notably exygen and water vapour, leaving an oxide ceramic layer with increased porosity. The bath temperature is maintained constant at about 20°C. A constant current density of at least 1A/dm²: is maintained in the electrolytic bath until the voltage reaches a predetermined end value, consistent with the formation of an insulating layer. Under these conditions, one obtains typically about 1 µm of ceramic oxide coating per minute. Ceramic coating thickness up to about 100µm can be obtained in 60 minutes, depending on barrier forming metal type and alloy. The required current density to initiate the plasma process may be as high as 25A/dm² if the applied metallic layer is rough and porous.

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The electrolytic plasma oxidation is preferably carried out in a weak aqueous alkaline electrolyte of pH in the range from 7 to 8.5, preferably in the range from 7.5 to 8, at temperatures of about 20°C, which means that the integrity of the

aubstrate material is little affected. As indicated above the melting that occurs during the formation of the ceramic coating tends to fill out any pores in the underlying metallic layer, resulting in an impermeable interfacial region between the layers.

For plastic substrates the formation of the ceramic oxide coating over the underlying metallic layer overcomes the problems of electrostatic repulsion commonly encountered when depositing ceramic particles directly onto the authorse of plastic substrates.

The substrate is preferably a component of a vacuum pump, and so the present invention also provides a vacuum pump component formed from metallic or plastics material and having a coating thereon formed by electrolytic plasma oxidation of a metallic layer applied to the component.

The invention will now be described with reference to the following non-limiting examples. Variations on these failing within the equips of the invention will be apparent to a person skilled in the art.

EXAMPLES

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In the present invention, one can achieve adherent and coherent ceramic coatings on iron castings, steels and plastics in a relatively simple and cost-effective manner that will also allow its application to precision parts with tight tolerances. The general method for achieving this is set out below, specific examples of which are presented thereafter.

(1.) An optional initial treatment to roughen the surface of the substrate. Such methods may include peening and blasting, pickling and/or combinations thereof. For plastics, application of a thin layer of liquid adhesive, such as polyimides or epoxies, or metal such as nickel, may follow the surface roughening.

- (2.) Deposition of a light metal (e.g., Al, Tl, Mg, and their alloys) or alloy (Al-Ni, Al-Cu, Al-Zn, Al-Mg, etc) onto the roughened surface (which may include a thin layer of liquid adhesive or metal), using . techniques such as sifting or compression of the metal powder or wrapping of the metal foll onto applied adhesive layer, or electrodeposition of the metal on to an initially-applied metal layer, vacuumplasma-spraying, sputtering. arc-epraying, spraying, high-valocity-oxy-fuel-spraying, and combinations thereof. In the case of a plactic substate, the much promising scatting techniques are the compression of the mutal powder or wrapping of the metal foil onto an applied liquid adhesive layer or electrodeposition of the metal on to initially-applied metal layer, the plasma spraying, the high velocity oxy-fuel spraying and combinations thereof, as these exhibit a low thermomochanical load with respect to other technologies. It will be appreciated that the above-mentioned My or a record to the tester as cored elitil event easign det private substrate.
- (3.) Electrolytic plasma oxidation of the surface of the metallic layer to generate a ceramic oxide coating. It is important that not all of the metallic layer is converted to ceramic.
- (4.) Optional finishing treatment of the surface of the ceramic coating using techniques such as keying in of substances (for example, CF_x, fluorocarbons, PTFE, MoS₂ and graphite, Ni, Cr, Mo, W and their Carbides, paints and resins), grinding, polishing, tumbling, rumbling, etc and combinations thereof.

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Example 1

A composite tube, manufactured in epoxy resin comprising carbon fibres (fibre direction to satisfy thermo-mechanical strain matching with metallic rotor parts), was subjected to the coating process. The surface of the tube was subjected to a low pressure grit blast using 60 mesh grit or light peening using baudte. Thermal sandblasting may also be used. All methods serve to remove the sheen from the surface of the tube, thereby to roughen the surface without damaging the fibres. The surface was then wiped with alcohol and dided to remove greace therefrom.

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Aleximism and aluminium-nickel alloy (20/20) having powders of nominal cizs ~10 μm were plasma sprayed onto the tube using a standard Ar/H₂ plasma, nominally of 40 kV/ power level. It is to be noted that use of standard powders with nominal dimension 45 - 90 μm tend to give a more porous coat. Each powder type resided for about 0.7 ms in the plasma at ~18000°C belong being projected and the tube, revolving at 60 pm, from a distance of 150 to 100 mm. The speed of the particles ampinging on the tube was in tange from 225 kV = 000 kVs, thus permitting splaying out (or wetting) of the molten particles and with some degree of penetration into the tube. The average surface temperature during the plasma spraying process was in the range 100 - 150°C. The coating thickness was controlled by the duration of the spraying. Following the spraying, the tube was slowly cooled in still air and the surface machined by grinding with a 180 SIC grinding wheel to remove the surface roughness, leaving a final thickness of the metallic layer thus formed on the tube of about 50 μm.

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The metallic layer applied as described above was subject to electrolytic plasma oxidation in an electrolyte (an aqueous solution of an alkali metal hydroxide and sodium silicate or sodium aluminate, or sodium metaphosphate), at a pH of 7.6. Using a current density of 12 A/dm²; an electrolyte temperature of 20±3°C, and a coating time of 60 minutes, a voltage end value of 350V was registered. The component with the thus-formed ceramic coating was washed and dried. The thickness of the ceramic coating was 30 µm.

The corrosion resistance of the composite tube coated in this manner has four times better corrosion resistance than un-coated epoxy-carbon fibre composite tube in semiconductor applications. In particular it was found that a BOC Edwards IPX pump having components coated with the ceramic coating lasted four times longer than un-coated pumps when exposed to 4500 litres each of chlorine, bromine and fluorine.

As a final, optional treatment, the coramic-social companent was immerced and moved within an equeous anionic PTTE dispersion having a publicle size of ~ 0.0 μm_e washed under a flow of hot water (00°C) and dried with hot air to entrance the convolon resistance of the coating.

Example 2

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A similar composite tube of example 1 was subjected to a low pressure grit black uping 20 mesh grit to remove the plant. Irom the surface of the pumper the plant to roughen the surface without damaging the fibres. The outfuse was then wiped with alcohol and dried to remove grease therefrom, prior to application of a thin liquid layer of epoxy adhesive using a paintbrush.

Aluminium and aluminium-nickel alloy (80/20) having powders of nominal size ~10 μm were compressed onto the surface of the tube by rolling compaction over a bed of the metal powder. Cure of the adhesive was achieved by placing the powder-coated tube for 1 hour in an oven pre-set to 120°C. The coating had an inner layer where the metal powder was intermixed with the adhesive and an outer layer where the powder was keyed onto the inner layer. Then the surface were machined by grinding with a 180 SIC grinding wheel to remove the surface roughness, leaving a final ground metallic layer thickness of about 30 μm.

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The metallic layer applied as described above was subject to electrolytic plasma oxidation in an electrolyte, (an aqueous solution of an alkali metal hydroxide and

sodium silicate or sodium aluminate, or sodium metaphosphate), at a pH of 7.6. Using a current density of 20 A/dm²; an electrolyte temperature of 20±3°C, and a coating time of 75 minutes, a voltage end value of 400V was registered. The tube with the thus-formed ceramic coating was washed and dried. The thickness of the ceramic coating was 10 µm. The corrosion resistance of the composite tube coated in this manner has four times better corrosion resistance than un-coated epoxy-carbon fibre composite tube in semiconductor applications.

The ceramic-coated tube can be optionally coated to enhance the corrosion resistance of the coating as in example :.

Example 5

Samples from example 2 above, with the ground metallic layer only, were further subjected to plasma spraying of aluminium and aluminium alloy powders under the conditions used in example 1. Following apraying, the tube was slowly cooled in will air. Then the surfaces were muchined by grinding with a 100 DIO untiding which is remove the surface roughness, leaving a final ground motallic layer thickness of about 30 µm.

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The metallic layer applied as described above was subject to electrolytic plasma oxidation in an electrolyte, (an aqueous solution of an alkali metal hydroxide and sodium silicate or sodium aluminate, or sodium metaphosphate), at a pH of 7.6. Using a current density of 12 A/dm²; an electrolyte temperature of 20±3°C, and a coating time of 60 minutes, a voltage end value of 350V was registered. The tube with the thus-formed ceramic coating was washed and dried. The thickness of the ceramic coating was 40 µm. The corrosion resistance of the composite tube coated in this manner has four times better corrosion resistance than un-coated epoxy-carbon fibre composite tube in semiconductor applications.

The ceramic-coated tube can be optionally coated to enhance the corrosion resistance of the coating as in example 1.

Example 4

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A similar composite tube of example 1 was subjected to a low pressure grit blast using 50 mesh grit to remove the sheen from the surface of the composite, thereby to roughen the surface without damaging the fibres. The surface was then wiped with alcohol and dried to remove grease therefrom, prior to application of a thin liquid layer of spory adhesive using a paintbrush.

An aluminium foil with a thickness of ~50 µm was wrapped onto the liquid adhesive. The outer diameter of the tube was coated by press rolling the tube over a cut section of the foil, and with the excess trimmed off, leaving an overlap length of ~1 mm. For the inner diameter, a similar out section of the foil was gontly laid around the surface, followed by consolidation with a roller, and with the access trimmed off, leaving an avendap length of ~1 mm. The office all solve was achieved by placing the foil-ocated tube for 1 hour in an oven proport to 120°C.

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The metallic layer applied as described above was subject to electrolytic plasma oxidation in an electrolyte, (an aqueous solution of an alkali metal hydroxide and sodium silicate or sodium aluminate, or sodium metaphosphate), at a pH of 7.6. Using a current density of 6 A/dm 2 ; an electrolyte temperature of 20±3°C, and a coating time of 45 minutes, a voltage end value of 300V was registered. The tube was subsequently washed and dried. The thickness of the ceramic coating formed on the tube was 35 μ m. The corrosion resistance of the composite tube coated in this manner has four times better corrosion resistance than un-coated epoxycarbon fibre composite tube in semiconductor applications.

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The ceramic-coated tube can be optionally coated to enhance the corrosion resistance of the coating as in example 1.

Example 5

A similar composite tube of example 1 was cleaned and the surface modified by roughening and activation, using grit blasting or its combination with plasma etching.

The modified polymer surface was then activated by Pd/Sn colloids to provide sites for deposition of a nickel layer by means of electroless nickel plating. A galvanic process that permits deposition of an aluminium layer onto the nickel layer (serving as bond coat) then follows. The typical coating thickness for the nickel layer was in the range from 5 to 23 jun, and the thickness of the overcoat aluminium layer was in the range from 15 to 50 µm. The coating to obtained was very adherent to the composite tube, smooth, non-porous and impairmeable to fluids.

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The metallic layer applied as described above was subject to electrolytic placina suidation in an electrolyte, (an aquasus solution of an alicali metal lightwoods and sodium silicate or sodium aluminate, or sodium metaphosphate), at a pli of 7.8. Using a current density of 4 A/dm2; an electrolyte temperature of 20±30C, and a coating time of 10 minutes, a voltage end value of 350V was registered. The tube was subsequently washed and dried. The thickness of the ceramic coating formed on the tube was 15 µm. The corrosion resistance of the composite tube coated in this manner has six times better corrosion resistance than un-coated epoxycarbon fibre composite tube in semiconductor applications.

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The ceramic-coated tube can be optionally coated to enhance the corrosion resistance of the coating as in example 1.

Example 6

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In this example, a SG iron sample, 100 mm x 100 mm x 5 mm, and a mild steel sample, 100 mm x 100 mm x 5 mm, were subjected to the coating process. The

surfaces of the samples were roughened by sandblasting, followed by a pickling in a 10% HF aqueous solution at room temperature for 60 minutes. The samples were then washed and dried.

The samples were then subject to plasma spraying of aluminium and aluminium alloy powders under the conditions used in example 1. Following spraying, the samples were slowly cooled in still air. Then the surfaces were machined by grinding with a 180 SiC grinding whost to remove the surface roughness, leaving a final ground metallic layer thickness of about 50 µm.

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The metallic layers applied as described above were subjected to elasticity to plasma oxidation in an electrolyte (an equecus solution of an alkali metal hydroxide and sodium silicate or sodium aluminate, or sodium metaphosphate) with a pH of 7.6. Using a current density of ~8 Mdm², an electrolyte temperature of 20±6°D and a couling time of 30 minutes, a voltage and value of 300V was registered. The samples were washed and dried. The thickness of the coramic socialing formed on the samples was ~50 pm. SB iron coaled in this manner has four times better conveinn resistance than descoaled SB from in semiconductor applications.

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The ceramic-coated samples can be optionally coated to enhance the corresion resistance of the coating as in example 1.

CLAIMS

- 1. A method of forming a coating on a metallic or plastics substrate, the method comprising the steps of applying a metallic layer to the substrate and forming the coating from the metallic layer by subjecting the metallic layer to electrolytic plasma oxidation.
- A method according to Claim 1, wherein the metallic layer is formed from one of aluminium, magnesium, titanium, tanialum, zirconium, neobydium, hainium, tin, tungsten, majybdanum, vanadium, antimony, bismuth, and alloys of the aforementioned metals.
- 3. A method according to Claim 1 or Claim 2, wherein the metallic layer is deposited on the substrate.
 - A mathod according to Glaim 3, wherein the metallic layer is agrayed on the substrate.
- 20 5. A method according to Claim 1 or Claim 2, wherein the metallic layer is adhered to the substrate.
 - A method according to any preceding claim, wherein the thickness of the metallic layer applied to the substrate is less than 100µm.
 - 7. A method according to any preceding claim, wherein the substrate is roughened prior to the application of the metallic layer thereto.

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8. A method according to any of Claims 1 to 6, wherein the metallic layer is formed on a second metallic layer previously applied to the substrate.

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9. A method according to any of Claims 1 to 6, wherein the metallic layer is formed on a second polymeric layer previously applied to the substrate. A method according to any preceding claim, wherein the substrate 10. comprises iron. 11. A method according to any of Claims 1 to 9, wherein the substrate is an epoxy-carbon fibre composite or fibre reinforced plastics material. 12. A method according to any preceding claim, whorsin the metallic layer is smoothened prior to the formation of the coating therefrom. A method according to any preceding claim, whorsin the electrolytic 13. plasma exidation is performed at a pH in the range from T to 0.5. 11 A method according to any preceding claim, wherein the thickness of the coating formed from the metallic layer is less than 100µm. A method according to Claim 19, wherein the thickness of the coating 15. formed from the metallic layer is less than 50µm. 16. A method according to any preceding claim, wherein the external surface of the coating formed from the metallic layer is subsequently treated to modify the physical and/or chemical properties of the coating formed on the substrate, 17. A method according to Claim 16, wherein an external layer of the

coating is at least partially removed following formation thereof from

the metallic layer.

- A method according to Claim 17, wherein at least part of the external layer is abrasively removed from the coating.
- 19. A method according to any of Claims 16 to 18, comprising applying to the coating a material for reducing the porceity of the coating.
 - 20. A method according to any of Claims 16 to 18, comprising applying to the ocating a material for subanding the continuous resistance of the coating.
- 21. A method according to any of Claims 13 to 20, comprising applying to the coating a layer formed from one of a fluorocarbon, polytetrafluoroethylene, MoSo, Carbon, Ni, Cr, Mo, W, carbides of any of the aforementioned metals, a paint and a resin.

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- A method according to any preceding claim, wherein the substrate is a component of a vacuum pump.
- A vacuum pump component formed from metallic or plastics material and having a coating thereon formed by electrolytic plasma oxidation of a metallic layer applied to the component.

ABSTRACT

A method of forming a coating on a metallic or plastics component of a vacuum pump comprises the steps of applying a metallic layer to the component and forming the coating from the metallic layer by subjecting the metallic layer to electrolytic plasma oxidation.

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